## PATENT SPECIFICATION

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## (54) EMULSIONS

(71) We, L'OREAL, a French Body Corporate of 14 Rue Royale 75, Paris 8e, France, do hereby declare the invention for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:-

The present invention relates to emulsions of the "water-in-oil" and "oil-in-water"

It has already been proposed to produce cosmetic products such as make-up or beauty creams which are in the form of "water-in-oil" emulsions because the water beauty creams which are in the solly phase ensures, in certain cases, better 10 moisturisation of the epidermis and better protection of the latter. However, it has hitherto been difficult to market cosmetic products of this type because such "water-

mentro ocea camera, to market comments products or uns type occasio such water-in-oil? enablishing sperrally display two sorts of difficulties:

Text of all, it is necessary that the emulsions should not invert, i.e. they should not convert into an emulsion of the "oil-in-water" type by dilution with water.

The products should be sufficiently studied to retain their finely dispersed state, regardless of the period of storage (which studies to retain their finely dispersed state, regardless of the period of storage (which 15 can be several years), and regardless of major variations in temperature which generally favour destruction of the emulsion through loss of the aqueous phase from the dispersed state, which furthermore is a particular hazard where the emulsions are

subjected to low temperatures. With this in mind, the Applicant Company has already proposed to use, as the emulsifier in such emulsions, a mixture of an oxypropylenated-polyglycerolated alcohol and of magnesium isostearate, succinate esters of polyoxyalkylenated fatty alcohols, or

oxypropylenated-oxyethyleneated alcohols.

Furthermore, it is already known to use, as emulsifiers in cosmetic compositions, in particular creams, polymers consisting of a sequence obtained by polymerisation of propylene oxide, to which have been grafted two sequences obtained by polymerisation of ethylene oxide. Such copolymers are known under the name of Pluronics

(sold by Messrs. Wyandotte Chem. Corp.). are known unuer the name or rutomes. The Applicant Company has now found, surprisingly, that it is possible to produce very good cosmetic canulsions by using a large variety of certain types of sequence polymers as the emulsifier.

Such polymers are known and some of them have already been proposed as additives in motor lubricants.

It is well known that the various monomers involved in the production of

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It is well known that the various monomers involved in the production or copolymers on behave in different ways in the formation of the polymer chain, and in general polymerisation processes make it possible to alant the reaction towards and the process in the possible to alant the reaction towards merisation process it is possible to obtain copolymers wherein the monomer units are grouped according to types, these groupings being described by the name of "sequence". Such copolymers are defined herein as "sequence copolymers". The sequence polymers are generally binary polymers containing two types of sequence, each made up from identical monomers. The number of sequences is sequence, each made up from identical monomers. The number of sequences

generally two or three.



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The copolymers containing two sequences are called "bisequence" copolymers, and the distribution of the monomer units in the polymer chain can be represented as follows:

The copolymers containing three sequences are generally called "trisequence" copolymers and the distribution of the monomer units in the polymer chain can be represented as follows:

The present invention provides an emolsion which can be used in cosmetics, of the "start-rin-til" type, which is stable and cannot readily be inverted; it contains as the emulsifier, a sequence opplymer simultaneously containing at least one lipophilis expenses and at least one hydrophilis expenses. The lipophilis esquences are obtained from monomers with lipophilis chains, whilst the hydrophilis expenses are obtained from monomers with hydrophilis chains. The lipophilis expenses at the sequence polymers used in the emulsions of this invention can be represented by the following formula: 10

R is selected from the group consisting of

each of R1 and R2, which may be identical or different, represents either a hydrogen atom or an alkyl radical with 1 to 4 carbon atoms, R<sub>2</sub> represents a saturated hydrocarbon chain of 6 to 18 carbon atoms,

R<sub>s</sub> represents a sammaten nyurocaron cream of 0 to 10 carona counts.

R<sub>s</sub> represents a methyl or ethyl radical and
R<sub>s</sub> represents a sammated hydrocarbon chain of 5 to 26 carbon atoms.

The hydrophilic sequences of the sequence polymers used in the emulsions of this invention can be represented by the following formula:

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in which:

R" is selected from the group consisting of:

(b) 
$$-C00-Y-N < R_{a'}$$

(c) 
$$-C00-Y-N<\frac{{R_1}'}{{R_2}'}$$
. HX

(d) —O≔N

(k) 
$$-\sqrt{R_1^{l'}}$$
 HX

R'" representing a methyl radical

R'" representing a hydrogen atom

- each of R<sub>1</sub>' and R<sub>2</sub>', which may be identical or different, represents either a hydrogen atom or an alkyl radical with 1 to 4 carbon atoms, Y represents a saturated hydrocarbon chain of 2 to 4 carbon atoms or a hydrocarbon chain of 2 to 4 carbon atoms interrupted by hetero-atoms such as oxygen and sulphur, and
  - HX represents an inorganic or organic acid taken from the group consisting of hydrochloric acid, hydrobromic acid, lactic and acetic acid. If R" represents a carboxylic acid group, this group can be neutralised with an
  - hydrocumic acid, phydrocramic acid, acute and acenc acid.

    If R" represents a carboxylic acid group, this group can be neutralised with an inorganic or organic base, such as ammonia, monoethanolamine, diethanolamine, triethanolamine, the isopropylamines, morpholine, 2-amino-2-methyl-propanol-1, or 2-

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ne monument winen can tean to the romaton of nyurophinic sequences, the following may be quoted: 2-vinyl-pyridine, its hydrochloride and its lactate; 4-vinyl-pyridine, its hydrochloride and its lactate; para-dimethylamino-styrene, its hydrochloride and its lactate; 2-(N,N-dimethylamino)-ethyl methacrylate; 2-(N,N-dimethyla in incarcy and in incarcy act, 2-(N,N-dimensylamino)-ethyl methacrylate, 2-(N,N-dimensylamino)-ethyl methacrylate, 2-(N,N-dimensylamino)-ethyl-glycol methacrylate, 2-(N,N-dimensylamino)-ethyl-glycol methacrylate; methacryloutrile.

In an alternative embodiment, the hydrophilic sequences containing the tertiary amine groups are quaternised by means of a quaternising agent chosen, for example, from dimethyl sulphate, ethyl bromide or  $\beta$ -bromoethanol.

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The molecular weights of the sequence polymers used according to the invention can vary within wide limits. They are generally fixed as a function of the properties

desired of the emulsifier. The sequence polymers according to the present invention generally have a molecular weight of between 1,000 and 1,000,000, but preferably between 8,000 and

700,000. Equally, the ratio of the lengths of the sequences can vary within very wide limits and is generally decided by the application for which the copolymer is destined, namely either a "water-in-oil" emulsion or "oil-in-water" emulsion.

In effect, the sequence polymers used according to the present invention can also be used to produce "oil-in-water" emulsions if the sequence polymer is soluble in water whilst having a certain affinity towards oils. One of the most characteristic and the most important properties of the sequence

of the of the most characteristic and the most important properties of the soughtee polymers is that each of the sequences displays the properties of the corresponding homopolymer. Depending on the choice of the sequences, it is possible to obtain sequence copolymers which are simultaneously hyrophilic and lipophilic, but of which the hydrophilic or lipophilic character is more or less marked.

The present invention also provides a cosmetic composition or an excipient for a 30

pharmaceutical product, which comprises an emulsion of the present invention. The proportion of emulsifier in the emulsion according to the invention can vary within very wide limits, for example from 5 to 20% by weight, whilst the proportion

of water can vary from about 20 to 75% by weight relative to the total weight of the 35 constituents. In general, the proportion of emulsifier relative to the mixture of oil+wax is at

least 10% by weight. The proportion of the mixture of oil+wax relative to the total weight of the emulsion according to the invention is generally between 20 and 65% by weight. According to the invention, a large variety of products can be used to form the

oily phase of the emulsions, such as:
hydrocarbon oils, such as paraffin oil, stringy "Vaseline" (Registered Trade Mark, i.e. partially flowing paraffin), perhydrosqualene and solutions of microcrystalline wax

in paraffin oil and Purcellin oil. animal or vegetable oils, such as caballine oil, pork fat, sweet almond oil, callo-hylum oil, olive oil and avocado oil, these being oils which are well absorbed by the skin but which can in certain cases give rise to rancidity;

sun out winch can in cream cases give use to function;
saturated esters which cannot turn rancial and have good penetrating ability, such
as isopropyl palmiatte, isopropyl myristate, ethyl palmiatte, diisopropyl adipate and
the triglycenides of octanoic and decanoic acids.
Silicone oils which are soluble in other oils or phenyl-ethyl alcohol can also be

added to the oily phase In certain cases it is possible to utilise waxes such as carnauba wax, candellila wax, beeswax, microcrystalline wax and ozokcrite to assist retention of the oils.

Long-chain fatty alcohols, such as the fatty alcohol from beeswax, cholesterol, Long-cnam may accounts, such as the lady account from Decimar, consistently, funding alcohol or magnesium startant can also be used as adjuvants to the oily phase. The emulsions according to the invention make it possible to produce the most reporture, such as moisturising creams, foundation creams, make-up, fluid creams, brilliantines and products for protection against sunburn.

A further subject of the present invention is a process for the preparation of "water-in-oil" and "oil-in-water" emulsions from emulsifiers consisting of the sequence

polymers defined above.

This process of preparation is essentially characterised by the fact that in a first stage the sequence copolymer is mixed with the "oil" phase with vigorous stirring and

	at a temperature of about 150°C and that, in a second stage, after having cooled the sequence copolymer-toil/wax mixture to a temperature of about 80°C, the "water" phase, with or without the addition of hydrochloric, lactic or acetic acid, and pre-	
5	viously heated to the same temperature, is introduced into it, after which the mixture is cooled to ambient temperature, whilst stirring. At the end of the operation, the emulsion can be passed through a (triple) roll mill to refine it.	5
	Though the process for the preparation of the sequence polymers is known in general we will review the principal stages involved.  These polymerisations are generally initiated by so-called "anionic" initiators,	
10	which are generally metals belonging to the first group of the periodic table of the elements, such as lithium, sodium and porassium, or organic compounds of these metals. Compounds such as diplenyl-methyl-sodium, fluorenyl-lithium, fluorenyl-lithium, fluorenyl-methyl-sodium, fluorenyl-lithium, fluorenyl-methyl-sodium, fluorenyl-lithium, fluorenyl-methyl-sodium, fluorenyl-lithium, fluorenyl-methyl-sodium, fluorenyl-methyl-methyl-sodium, fluorenyl-methyl-methyl-sodium, fluorenyl-methyl-methyl-sodium, fluorenyl-methyl-methyl-sodium, fluorenyl-methyl-methyl-sodium, fluorenyl-methyl-methyl-sodium, fluorenyl-meth	10
15	sodium, naphthalene-sodium, naphthalene-potassium, naphthalene-thibium, tertaphenyl- disodiobutane and phenyl-isopropyl-potassium may, for example, be mentioned. The choice of the polymerisation initiators is in fact very important, because it	
5	allows the structure of the sequence polymer to be decided. Thus, naphthalene-sedium allows the structure of the sequence polymer to be decided. Thus, naphthalene-sedium allows the polymerisation to be directed towards obtaining a "tri-sequence" copolymer. On the other hand, phenylisopropyl-potassium allows the polymerisation to be directed towards obtaining a "bi-sequence" polymer.	15
20	I nese polymerisation reactions leading to the formation of sequence polymers take place in aprotic solvents such as, for example, benzene, tetrahydrofurane and toluene.	20
25	In general terms, tri-sequence polymens, for example, are obtained as follows. First of all, a solution of the initiator in the selected objects is prepared, and then one of the monomers which is to give rise to one of the sequences is added; after polymerisation of this monomer (this polymerisation taking place in the space of a few minutes), the second monomer which is to give rise to the formation of the two other sequences is added, and these two other sequences is added, and these two other sequences is added, and these two other sequences.	25
30	metrically relative to the sequence of the first monomer. After the end of the poly- merisation, the tri-sequence polymer can be deactivated by means of a few drops of methanol.	30
35	In general, the reaction leading to the formation of these sequence polymers is carried out at a temperature of about -70°C. These polymerisation reactions aimed at the production of sequence polymers can obviously not be carried out with monomers containing mobile hydrogens, such as acids and amides. Hence, if it is desired to obtain sequence polymers containing acid or amide.	35
40	groups in one of their sequences, it is necessary to start from monomers which can subsequently, through chemical reaction, give rise to this type of group. For example, it is possible, for this purpose, to start from monomers possessing a nitrile group or	
40	an ester group. In effect, it is possible to obtain the corresponding acids by hydrolysis, and subsequently to obtain the corresponding amides by amidification.  Such a procedure can be used if it is desired to obtain lipophilic sequences consisting of methacylamide radicals or hydrophilic sequences consisting of methacylle aid radicals.	40
45	The following Examples further illustrate the present invention. Example 1 illustrates the preparation of the sequence polymers.	45
	EXAMPLE 1.  Preparation of a bisequence polymer of 2-vinylpyridine and lauryl methacrylate  One litre of anhydrous distilled tertahydrofurane is introduced into a flask of two	
50	litres capacity equipped with a mechanical stirrer, two dropping funnels, a graduated tube, a nitrogen inlet tube, a dip tube which allows samples of the reaction mixture to be taken during the reaction, and a thermometer. The flash is then cooled to a tem-	50
	perature of $-70$ °C by means of a mixture of solid carbon dioxide and methanol.	
55	The whole of the apparatus is under a nitrogen atmosphere, which apparatus has	-
22	been carefully purified by heating to 400°C in the presence of copper foil, and the stream of nitrogen is also purified by passing over anhydrous potassium hydroxide and over anhydrous magnesium perchlorate. A solution of diphealy-methyl-sodium in anhydrous distilled tetrahydrofurane is	55
	added dropwise by means of the graduated tube, whilst stirring. At the start of the	
60	addition, the diphenyl-methyl-sodium solution loses its colour as soon as it comes into contact with the tetrahydrofurane of the flask. The introduction of the diphenyl-methyl-sodium solution is then continued until a reddish-yellow colour persists in the	60
	reaction flask. A further 2.82 ml of a solution containing 247 mg of diphenyl-methyl-	

U				
	sodium in tetral	hydrofurane are then introduced through the	he same graduated tube, the	
	30 3 m of c	er a nitrogen atmosphere. arefully purified 2-vinylpyridine are intro of the dropping funnels, under nitrogen	duced rapidly into the flask	
5	The tempe	rature inside the flask rises to $-62^{\circ}$ C for ction mixture becomes darker.	or a few minutes, whilst the	5
	A amoll on	nount of solution of "living" polymer of 2- ed by suction, using the dip tube, and is	-vinylpyridine in tetrahydro- employed for calculating its	
10	molecular weigh	it.	again, 22.4 g of carefully	10
10	purified lauryl other dropping	methacrylate are rapidly introduced into funnel, under nitrogen. The temperature	rises to -62°C and when	
				10
15	deactivated. In methanol. The distilled, and th	general, this last stage is carried out of solution then becomes practically colourle are residual polymer dissolved in chlorofor	less, the tetrahydrofurane is m and then precipitated by form and twice precipitating	15
20		ether, the polymer is dried under reduce polymer (yield, 60%) are thus obtained rmined by the light staggering method in		20
		$\overline{M}_p = 110,000, d_n/d_c \text{ (MeOH)} = 0.$	.184.	
25	deactivated with purification of t	le of the homopolymer of poly-2-vinyl n methanol and purified in accordance wi he bisequence polymer, can be used to de		25
	in the same way	: $M_p = 60,000$ , $d_n/d_c$ (MeOH)=0.3	236.	
:	The seque accordance with	nce polymers shown in Tables I and I the procedure as described above.	II below were prepared in	•
30		EXAMPLES OF COMPOSITION	ONS	30
		EXAMPLE A:		
	A fluid cre	eam of the following composition is prep		
		Copolymer No. 3	7 g 40 g 3 g	35 -
35		Paraffin oil	40 g	
		Microcrystalline wax Water	7 g 40 g 3 g 50 g	
		EXAMPLE B:		
40	A foundati	on cream of the following composition is	prepared in accordance with	40
.40	the invention.	Copolymer according to Example 1 Paraffin oil	7.4 g 20 g	
		Perhydrosqualene	24 g	
		Titanium oxide	1.5 g	45
45		Titanium oxide Ochre	1.5 g	45
45		Titanium oxide Ochre Períume	1.5 g 0.2 g	45
45		Titanium oxide Ochre	1.5 g	45
45	A night cre	Titanium oxide Ochre Períume	1.5 g 0.2 g 45.4 g	
<b>45</b>	A night cre	Titanium oxide Ochre Perfume Water.+lactic acid (3.4 g)  EXAMPLE C: am of the following composition is prepare Copolymer according to Example 1	1.5 g 0.2 g 45.4 g d according to the invention:	45
	A night cre	Titunium oxide Ochre Perfume Water+lactic acid (3.4 g)  EXAMPLE C: am of the following composition is prepare Copolymer according to Example 1 Parafin oil	1.5 g 0.2 g 45.4 g d according to the invention:	
	A night cre	Titunium oxide Ochre Perfume Water+lactic acid (3.4 g)  EXAMPLE C: am of the following composition is prepare Copolymer according to Example 1 Paraffin oil Isopropyl palmitate Purcellia oil	1.5 g 0.2 g 45.4 g  d according to the invention: 7 g 22.1 g 10 g 12 g	50
	A night cre	Timium oxide Ochre Perfume Water+lactic acid (3.4 g)  EXAMPLE C: am of the following composition is prepare Copolymer according to Example 1 Paraffin oil Isopropyl palmitate	1.5 g 0.2 g 45.4 g d according to the invention: 7 g 22.1 g 10 g	

"Bisequence" Polymer TABLE I

L/H in weight in the	66/34	90/10	90/10	91/9	31/69	50/50
Elementary analysis	C 73 H 9.6 N 4.5	C 74.4 H 11.6 N 0.9	C 74.5 H 11.4 N 0.9	C 74.5 H 11.4 N 0.8	C 87.9 H 7.6 N 4.1	C 85.1 H 7.5 N 6.6
4   8E	0.113	0.079	0.079	0.079	0.181	0.182
Average molecular weight	967,000	8,000	109,000	254,000	15,400	270,000
Yield	45	14	15.8	32	19	8
Amount of catalyst	"	4,400(a)	430(a)	430(a)	1,260(b)	1,260(b)
Catalyst solution in THF	8	93		v	22	52
Amount	15	12.5	12.5	27.5	5	01
. Am	98	2 .	7	7	7	7
Monomer 2	Lauroyl methacrylate (L)	£3.	ĘĴ.	<b>£</b>	2-Vinyl- pyridine (H)	£
Monomer 1	2-Vinyl- pyridine (H)	2-Dimethyl- amino-ethyl methacrylate (H)	Ħ	Ħ	Styrene (L)	Ð
Copolymer	2	6	4	10	9	

N.B.: The letter "T." signifies lipophilic The letter "T" signifies hydrophil (a) Diphenyl-methyl-sodium (b) Phenyl-isopropyl -potassium

The preparation of the above "tistegrance" copolymens is carried out by homopolymentsation of the monomer 1, followed by copolymentsation of homopolymer with the monomer 2.

"Trisequence"

		ī				
Copolymer No.	Monomer 1	Monomer 2	Type of copolymer	Quar 1	2	Catalyst solution in THF (ml)
8	Styrene (L)	2-Vinyl pyridine (H)	н—нц—цн—н	20	5	12
9	" (L)	" (H)	H—HL—LH—H	20	13	12
10	" (L)	" (H)	н—ні.—і.н—н	20	30	12
11	4-Methyl styrene (L)	4-Vinyl- pyridine (H)	н-нг-гн-н	. 6	1.5	. 3
12	" (L)	" (H)	H-HL-LH-H	6	4	3
13	Styrene (L)	2-Dimethyl amino-ethyl methacrylate (H)	н—ні.—і.н—н	20	5	12
14	" (L)	" (H)	H-HL-LH-H	20	13	12
15	" (L)	" (H)	H-HL-LH-H	20	30	12
16	2-Vinyl- pyridine (H)	Lauryl methacrylate (L)	L—LH—HL—L	11	5	* .
17	, (H)	" (L)	L-LH-HL-L	- 11	11	6
18	Lauryl methacrylate (L)	2-Dimethyl- amino-ethyl methacrylate (H)	н—ні—ін—н	5	2	6
19	" (L)	" (H)	H-HL-LH-H	5	5	6
20	Styrene (L)	4-Vinyl- pyridine (H)	H—HL—LH—H	5	3	. 3
21	" (L)	" (H)	H-HL-LH-H	.5	10	. 3
22	" (L)	2-Dimethyl- aminoethyl methacrylate	н-нг-гн-н	20	2	12
		(H)		<u> </u>	·	erication of th

N.B. The preparation of the acove "trisequence" copolymers is carried out by homopolymerisation of the monomer I, followed by copolymerisation of the homopolymer with the monomer 2. "L" denotes "lipophilic" and "H" denotes "hydrophilic".

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## Copolymer

Amount of catalyst (mg) naphthalene	Yield	Average molecular	dn dc	Elementary analysis		L/H, by weight	
sodium	%	weight	(THF)	C	н	N	in the copolymer
543	80	52,000	0.182	91.3	9.0	<1	>93/7
543	76	57,000	0.174	84.1	. 7.3	7.7	42/58
543	74	50,000	0.178	89.5	7.8	2.4	82/18
408	40	55,000	0.183	91	8	<1	>93/7
408	35	71,000	0.167	89.7	8	1.3	89.9/9.1
543	60	78,000	0.153	82.5	8.5	2.6	71/29
				-		-	
543	55	196,000	0.142	77.3	8.5	2.6	58.5/41.5
543	50	202,000	0.131	74.5	7.0	4.5	49.5/50.5
816	50	40,000	0.150	76.4	9.5	5.3	61/39
816	41	46.000	0.116	76.4	9.7	5.4	60/40
816	20	730,000	0.082	58.9	9.2	6.8	27/75
						1	
816	40 .	880,000	0.080	61.4	9.8	7.4	17/83
407	44	66.000	0.189	92.2	7.5	<1	>92.5/7.5
407	27	65.008	0,195	92.7	7	<1	>92.5/7.5
543	60	78,000	0.153	82.5	8.5	2.6	71/29

10	1,324,745				10
	EXAMPLE D: A milk of the following composition is prepared acc	cording to	o the inve	ention:	
	Copolymer No. 4 Paraffin oil Stringy vascline	12 18 8	g g g		5
5	Triglyceride of octanoic and decanoic acid Ozokerite	10 2 50	g g		
10	Water + acetic acid  EXAMPLE E:	-	В		10
	A cheek make-up is prepared according to the inv	ention:			
15	Copolymer No. 10 Stringy vaseline 2-O-Cityl-dodecanol-1 Isopropyl palmitate Diisopropyl adipate Candellila wax	15 6 2 5 37.4 2.5 2	8 8 8 8 8 8 8		15
20	Carnauba wax D and C Red No. 8 (dyestuff)  CH  CH  CH  SO, Ha	-	5		20
	Red iron oxide	0.5	g		
	Titanium oxide Water	1.5 28	g .		
25	EXAMPLE F: A moisturising milk for protection against sunburn	n is prep	ared acco	rding to the	25
	invention:  Copolymer No. 11  Triglyceride of octanoic and	10	g		30
30	decanoic acid Isopropyl myristate Disopropyl adipate Oxologiie	6 11 30 2	8 8 8 8 8 8		50
35	"Parsol-Ultra" sold by Messrs. GIVAUDAN (a mixture of aminobenzoic acid esters and				35
	substituted cinnamic acid esters; filter for sunlight) Water+lactic acid (0.6 g)	39	g g		
40	EXAMPLE G: A tinted "open air" cream is prepared according	to the in	vention:		40
	Copolymer No. 13	10	g		
	Copolymer 140. 13  Isopropyl palmitate  Diisopropyl adipate  Paraffin oil	7 28	g g		45
45	Beeswax Red iron oxide Yellow iron oxide	6 2 1 1	g g g		
50	Titanium oxide Water + hydrochloric acid (0.7 g)	44	g		50

	EXAMPLE H: A cuticle cream is prepared according to the invention:	
5	Copolymer No. 16 7 g Isopropyl painitane 20 g Perhydrogundene 30 g Stringy vaseline 7 g CO 3 g 2-Octyl-dodecanol-1 3 g Water 30 g	5
10	EXAMPLE 1:  An "oil-in-water" make-up remover cream is prepared according to the invention:	10
15	Copolymer No. 18	15
	EXAMPLE 1: A cream of the following composition is prepared according to the invention:	
20	Capolymer No. 9   15 g   Phenyl-ether alcohol   40 g   Disorproyl adipate   7 g   Water+acetic acid (4 g)   38 g	20
25	EXAMPLE K: A cream of the following composition is prepared according to the invention:	25
	Copolymer No. 6	
30	Stringy vaseline 1 g Disopropyl adipate 26 g Ozokerite 2 g Water 42 g	30
	EXAMPLE 1.  A cream of the following composition is prepared according to the invention:	
35	Copolymer No. 3 10 g Perhydrosqualene 25 g Stringy vaseline 14.5 g Ozokerite 3 g	35
	Water 47.5 g	
40	The emulsions according to the invention are particularly suitable for the preparation of foundation creams, make-up and hand creams.  Of course the embodiments of the invention which have been described are given merely by way of illustration and numerous modifications are possible. In particular its ideat that it is possible to use several emulsifiers according to the invention in the invention of the invention in the invention of the invention in the inventi	40
45	It is clear that it is possible to the several enhancement according to the invention simultaneously, optionally together with other previously known emulsifiers. It is also obvious that all the ingredients usually employed, and in particular those like that only improve the sability and shelf life of the emulsions, can be introduced into the emulsions according to the invention. Finally, it will be understood that the emulsions according to the invention can also be used in fields other than those of	45
50	cosmetics and of excipients for pharmaceutical products.	50

WHAT WE CLAIM IS:

1. An emulsion of the "water-in-oil" or "oil-in-water" type which comprises, as emulsifier, at least one sequence polymer (as hereinbefore defined), which contains (i) at least one lipophilic sequence corresponding to the formula:

$$- \begin{bmatrix} c_{12} - c_{1}^{p'} - c_{12} - c_{2}^{p'} - c_{12} - c_{2}^{p'} - c_{12} - c_{2}^{p'} - c_{12} - c_{2}^{p'} \\ k & k & k \end{bmatrix}$$

in which:

R represents a radical of the formula:

in which case R' represents a hydrogen atom, or 10

in which case R' represents a methyl radical,
each of R, and Rs, which may be the same or different, represents a hydrogen
atom or an allyly radical with 10 4 carbon atoms,
R, represents a stantated hydrocarbon chain of 6 to 18 carbon atoms,
R, represents a methyl or ellyl radical, and
R, represents a testumed hydrocarbon chain of 5 to 26 carbon atoms,
and (ii) at least one hydrophilic sequence corresponding to the formula:

in which:

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R" represents a radical of the formula:

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in which case R" represents a methyl radical,

(e)	<b>-</b>
<b>(f)</b>	<b>-</b> ⊘
(g)	Xx . 10X
(h)	~ <u>\</u>
<b>(</b> )	-K. 11X
<b>(</b> )	$- \hspace{-1.5cm} \bigcirc \hspace{-1.5cm} \stackrel{R_1^I}{\sim} \hspace{-1.5cm} \stackrel{R_2^I}{\sim} -1.5$
or (k)	- $        -$

in which case R''' represents a hydrogen atom
each of R,' and R<sub>2</sub>, which may be the same or different, represents a hydrogen
atom or an alley radical with 1 to 4 carbon atoms,
Y represents a saturated hydrocarbon chain of 2 to 4 carbon atoms, optionally 10 10 containing one or more chain hetero-atoms, and

HX represents hydrochloric acid, hydrobromic acid, lactic acid or acetic acid.

2. An emulsion according to Claim 1 wherein Y represents a saturated hydrocarbon chain of 2 to 4 carbon atoms containing at least one chain oxygen or subplur 15 3. An emulsion according to Claim 1 or 2 wherein R" represents a carboxylic

An emusion according to Chim I or Z wherein N represents a caronayue-acid group which has been neutralised with an inorganic or organic base.

An emulsion according to Chim I or Z wherein R" represents a carboxylic said group which has been satisfied by a sodium, possissim or magnesism sail.

An emulsion according to any one of the preceding claims wherein the hydro-lation of the control of the proceeding claims wherein the hydro-lation of the control of the preceding claims wherein the hydro-lation of the control of the

philic sequence is derived from 2-vinylpyridine, its hydrochloride or lacate, 4-vinyl-pyridine, its hydrochloride or lacate, para-dimenlylaminostyreae, its hydrochloride or lacate, 2-N.N-dinethylamino) ethyl methacrylate, 2-(N.N-dichylamino) ethyl methacrylate, 2-(N.N-dichylamino) ethyl-glyon methacrylate, 2-(N.N-dichylamino) ethyl-glyon methacrylate or methacrylanitie.

6. An emulsion according to one of the preceding Claims wherein the hydrophilic second of the control of the rised by dimethyl sulphate, ethyl brounds or gl-rownochtand.

8. An emulsion according to any one of the preceding claims wherein the lipo-nitic sequence is derived from styrene. 4-methalwayerse of lauryl methacrylan

30 philic sequence is derived from styrene, 4-methylstyrene or lauryl methacrylate.

	9. An emulsion according to any one of the preceding claims, wherein the	
	10 An emulsion according to Claim 9 wherein the sequence polymer has a	
	molecular weight of between 8,000 and 700,000.  11. An emulsion according to any one of the preceding claims wherein the	5
5		
	amount of oil, and wax if present, is between 20 and 65% by weight based on the	10
10		
	14. An emulsion according to any one of the preceding claims wherein the	
15		15
15	represent actor which does not turn rancid and is pencirating to the sain.	
20	myristate, ethyl palmitate, diisopropyl adipate or a triglyceride of octanoic or decanoic	20
20		
	17 As amulaion according to any one of the preceding claims wherein the '011	
	phase contains a carnauba wax, candellila wax, beeswax, microcrystalline wax or	
		25
25		25
23	19. A cosmetic composition which comprises an emulsion as claimed in any one	
		30
30		
	21. A composition according to claim 19 substantially as according to	
	Examples A to L.	
	as claimed in any one of claims 1 to 18 which comprises mixing the desired sequence	35
35	polymer with the "oil" part at a temperature of about 150°C, adding the "water"	
	part, previously heated to a temperature of about 80°C, to the mixture, heated to	
	about 80°C, with stirring, and cooling the mixture to ambient temperature with	
	stirring.  23. A process according to claim 22 wherein at least one of acetic acid, lactic	
	acid and hydrochloric acid is added to the "water" part.	40
40	24. A process according to claim 22 or 23 wherein the emulsion is subsequently	
	passed over a (triple) roll mill to refine it.  25. A process according to any one of claims 22 to 24 wherein the sequence	
	26. An emulsion whenever obtained by a process as claimed in any one of claims	45
45		
	22 to 25.	

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